10/574,098

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et fearl and fearl and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   18 -- 11
 chain nodes :
 2 3 5 10 11 13 14
 ring nodes :
 37 38
 ring/chain nodes :
 chain bonds :
 2-3 2-26 3-4 4-5 5-29 10-11 10-15 11-13 11-14 20-35 23-38
 ring bonds :
 15-16 \quad 15-20 \quad 16-17 \quad 17-18 \quad 18-19 \quad 19-20 \quad 22-23 \quad 22-27 \quad 23-24 \quad 24-25 \quad 25-26 \quad 26-27 \quad 25-26 \quad 25-26 \quad 26-27 \quad 25-26 \quad 25-2
 28-29 28-33 29-30 30-31 31-32 32-33 34-35 34-38 35-36 36-37 37-38
 exact/norm bonds :
 2-3 \quad 2-26 \quad 3-4 \quad 4-5 \quad 5-29 \quad 10-11 \quad 10-15 \quad 11-13 \quad 11-14 \quad 20-35 \quad 23-38 \quad 34-35 \quad 34-38
35-36 36-37 37-38
 normalized bonds :
 15-16 \quad 15-20 \quad 16-17 \quad 17-18 \quad 18-19 \quad 19-20 \quad 22-23 \quad 22-27 \quad 23-24 \quad 24-25 \quad 25-26 \quad 26-27 \quad 25-27 \quad 25-26 \quad 25-2
 28-29 28-33 29-30 30-31 31-32 32-33
 isolated ring systems :
 containing 15 : 22 : 34 :
 G1:Ak, O, S
```

G2:0,S

Match level: 2:CLASS 3:CLASS 4:CLASS 5:CLASS 10:CLASS 11:CLASS 13:CLASS 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom 20:Atom 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:Atom 28:Atom 29:Atom 30:Atom 31:Atom 32:Atom 33:Atom 34:Atom 35:Atom 36:Atom 37:Atom 38:Atom

=> s 16 sam

0 SEA SSS SAM L6

=> s 16 full

L8 1 SEA SSS FUL L6

=> dis

10/574,098

```
L8
    ANSWER 1 OF 1 REGISTRY COPYRIGHT 2009 ACS on STN
RN
     849675-84-9 REGISTRY
ΕD
     Entered STN: 03 May 2005
     4H-1-Benzopyran-2-carboxamide, N-[2-[2-[4-[2-[[2-(3,4-
CN
     dimethoxyphenyl)ethyl]methylamino]ethyl]phenyl]-2H-tetrazol-5-yl]-4,5-
     dimethoxyphenyl]-4-oxo- (CA INDEX NAME)
OTHER NAMES:
     4-Oxo-4H-chromene-2-carboxylic acid
CN
     N-[2-[2-[4-[2-[4-(2-(3,4-dimethoxyphenyl)ethyl]methylamino]ethyl]phenyl]-2H-
     tetrazol-5-yl]-4,5-dimethoxyphenyl]amide
     C38 H38 N6 O7
MF
SR
     CA
LC
     STN Files:
                CA, CAPLUS, CASREACT, TOXCENTER, USPATFULL
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PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> file caplus

=> s 18

L9 1 L8

=> dis 19 bib abs

L9 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2009 ACS on STN

AN 2005:324151 CAPLUS Full-text

DN 142:373847

TI Preparation of [(isoquinolinyl)ethyl]phenyl tetrazoles as P-glycoprotein inhibitors

IN Bang, Keuk Chan; Cha, Mi Young; Ahn, Young Gil; Ham, Young Jin; Kim, Maeng Sup; Lee, Gwan Sun

PA Hanmi Pharm. Co., Ltd., S. Korea

SO PCT Int. Appl., 53 pp. CODEN: PIXXD2

DT Patent LA English

FAN.	CNT 1 PATENT NO.							DATE		APPLICATION NO.						DATE		
ΡI	WO									WO 2004-KR2550						20041006		
		W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	ΒA,	BB	, BG	, BR,	BW,	BY,	BZ,	CA,	CH,
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ	, EC	, EE,	EG,	ES,	FI,	GB,	GD,
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS	, JP	, KE,	KG,	KP,	KΖ,	LC,	LK,
			LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK	, MN	, MW,	MX,	MZ,	NA,	NI,	NO,
			NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC	, SD	, SE,	SG,	SK,	SL,	SY,	ТJ,
			TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ	, VC	, VN,	YU,	ZA,	ZM,	ZW	
		RW:	BW,	GH,	GM,	ΚE,	LS,	MW,	MZ,	NA,	SD	, SL	, SZ,	TZ,	UG,	ZM,	ZW,	ΑM,
			ΑZ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM,	ΑT	, BE	, BG,	CH,	CY,	CZ,	DE,	DK,
			EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	ΙT	, LU	, MC,	NL,	PL,	PT,	RO,	SE,
			•			BF,	ВJ,	CF,	CG,	CI,	CM	I, GA	, GN,	GQ,	GW,	ML,	MR,	ΝE,
			SN,	TD,	ΤG													
	KR 2005033734 AU 2004277475				Α		20050413 20050414		KR 2003-69582									
									AU 2004-277475									
		2004277475						2007										
		2541301 1678162						20060712			CA 2004-2541301							
	ΕP									EP 2004-774778								
		R:	•				•	•	•	•		•	, LI,	•	NL,	SE,	MC,	PT,
			•	SI,	FΙ,	•						•	, PL,					
		CN 1863795									CN 2004-80029356							
	BR 2004015053 JP 2007507493 RU 2317985					A		20061128										
	JР	2007		Τ							-532094			20041006				
	RU	2317		C1								-114427						
	US	S 20070072900								US 2006-574098								
		MX 2006PA03803 IN 2006DN02327						20060703										
								20070713				2006-DN2327						
		NO 2006002019				А	20060706			NO 2006-2019					20060505			
PRAI		2003		82		A		2003										
		2004		550		W		2004		0.45								
OS GI	CAS	SREAC	Т 14	2 : 37	3847	; MA	RPAT	: 142	:373	847								

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- AB Title compds. represented by the formula I [wherein R1 = (un)substituted (hetero)aryl, acryl(hetero)aryl, heterocycloalkenyl, carbocyclo; R2-R11 = independently H, OH, halo, nitro, alkyl, alkoxy or R6R11 = cyclic ring; m, n = 1-4; X = CH2, O or S; and pharmaceutically acceptable salts thereof] were prepared as P-glycoprotein inhibitors. For example, II was given in a multistep synthesis starting from the reaction of 4-nitrophenyl bromide with 6,7-dimethoxy-1,2,3,4-tetrahydroisoquinoline•HCl. I showed inhibition of P-glycoprotein measured by using MCF-7 and MCF-7/Dx cell line and were tested for in vivo absorption of orally administered paclitaxel. Thus, I and their pharmaceutical compns. are useful as P-glycoprotein inhibitors for the treatment of cancers.
- RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

^{=&}gt; log y STN INTERNATIONAL LOGOFF AT 09:22:04 ON 14 JAN 2009